AMENDMENTS TO THE CLAIMS:

This listing of claims will replace all prior versions, and listings, of claims in the application:

 (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

an elongate composite structure adapted to externally constrict the stomach, said elongate composite structure comprising:

an elongate member for constricting the stomach of the patient.

a base material surrounding the elongate member, the base material making said composite structure self-supporting, and

a property improving means for improving at least one physical property of said composite structure other than self-supporting properties, said property improving means including at least one layer or coating applied on said base material, and

an adjustment means adapted to adjust the elongate member, and thereby, the composite structure to either enlarge or restrict the stoma opening,

the base material comprising a layer of polyurethane and a layer of silicone.

2. - 4. (Cancelled)

 (Withdrawn) An implantable constriction device according to claim 2, wherein said property improving means comprises a core of a viscoelastic material covered with said

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self-supporting base material.

6. (Withdrawn) An implantable constriction device according to claim 5, wherein

hard silicone constitutes said base material.

7. (Withdrawn) An implantable constriction device according to claim 5, wherein

said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel,

and collagen gel.

8. (Withdrawn) An implantable constriction device according to claim 2, wherein

said base material forms an inflatable tubing.

9. (Withdrawn) An implantable constriction device according to claim 8, wherein

said tubing has an inner surface defining the interior of said tubing, and said coating covers

said inner surface.

10. (Withdrawn) An implantable constriction device according to claim 8, wherein

said coating is selected from the group consisting of Tetrafluoroethylene polymer TeflonTM,

a poly-para-xylylene polymer Parylene™, and a biocompatible metal coating.

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11. (Withdrawn) An implantable constriction device according to claim 10, wherein

the biocompatible metal coating is selected from the group consisting of gold, silver and

titanium.

12. (Withdrawn) An implantable constriction device according to claim 8, wherein

hard silicone constitutes said base material.

13. (Withdrawn) An implantable constriction device according to claim 8, wherein

said base material forms two coaxial tubular layers and said property improving means

comprises a tubular intermediate layer of a viscoelastic material located between said coaxial

tubular layers.

14. (Withdrawn) An implantable constriction device according to claim 13, wherein

said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel,

and collagen gel.

15. (Withdrawn) An implantable constriction device according to claim 8, wherein

said base material forms an outer tubular layer, an inner arcuate layer attached to said outer

tubular layer, said outer and inner layers defining a curved space extending longitudinally

along said tubing, and said property improving means comprises viscoelastic material filling

said space.

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16. (Withdrawn) An implantable constriction device according to claim 15, wherein

said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel,

and collagen gel.

17. (Currently Amended) The implantable constriction device according to claim 1,

wherein said layer of said property improving means comprises a layer or coating on said

base material at least along a side of said elongate composite structure that is intended to

contact the stomach.

18. (Currently Amended) The implantable constriction device according to claim 17,

wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene

polymer, a poly-para-xylylene polymer, and a biocompatible metal coating.

19. (Currently Amended) The implantable constriction device according to claim 18,

wherein the biocompatible metal layer or coating is selected from the group consisting of

gold, silver and titanium.

20. (Withdrawn) An implantable constriction device according to claim 17, wherein

said property improving means comprises a core of a viscoelastic material covered with said

self-supporting base material.

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21. (Withdrawn) An implantable constriction device according to claim 20, wherein

hard silicone constitutes said base material.

22. (Withdrawn) An implantable constriction device according to claim 20, wherein

said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel,

and collagen gel.

23. (Withdrawn) An implantable constriction device according to claim 17, wherein

said base material forms an inflatable tubing.

24. (Withdrawn) An implantable constriction device according to claim 23, wherein

said tubing has an inner surface defining the interior of said tubing, and said coating covers

said inner surface.

25. (Withdrawn) An implantable constriction device according to claim 23, wherein

said coating is selected from the group consisting of a Tetrafluoroethylene polymer

TeflonTM, a poly-para-xylylene polymer ParyleneTM, and a biocompatible metal coating.

26. (Withdrawn) An implantable constriction device according to claim 25, wherein

the biocompatible metal coating is selected from the group consisting of gold, silver and

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titanium.

27. (Withdrawn) An implantable constriction device according to claim 23, wherein

hard silicone constitutes said base material.

28. (Withdrawn) An implantable constriction device according to claim 23, wherein

said base material forms two coaxial tubular layers and said property improving means

comprises a tubular intermediate layer of a viscoelastic material located between said coaxial

tubular layers.

29. (Withdrawn) An implantable constriction device according to claim 28, wherein

said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel,

and collagen gel.

30. (Withdrawn) An implantable constriction device according to claim 23, wherein

said base material forms an outer tubular layer, an inner arcuate layer attached to said outer

tubular layer, said outer and inner layers defining a curved space extending longitudinally

along said tubing, and said property improving means comprises a viscoelastic material

filling said space.

31. (Withdrawn) An implantable constriction device according to claim 30, wherein

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said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel, and collagen gel.

- 32. (Withdrawn) An implantable constriction device according to claim 1, wherein said base material forms a first layer and said property improving means comprises a second layer applied on said first layer, said second layer being more fatigue resistant than said first layer.
- 33. (Withdrawn) An implantable constriction device according to claim 32, wherein said second layer covers said first layer of said base material along a side of said elongate composite structure that is intended to contact the stomach.
- 34. (Withdrawn) An implantable constriction device according to claim 32, wherein said second layer comprises a polyurethane layer.
- 35. (Withdrawn) An implantable constriction device according to claim 32, wherein said property improving means comprises a coating coated on said first layer and/or said second layer, said coating having better aggressive body fluid resistance properties and/or better anti-friction properties than said base material.
 - 36. (Withdrawn) An implantable constriction device according to claim 35, wherein

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said coating is selected from the group consisting of Tetrafluoroethylene polymer $Teflon^{TM}$,

a poly-para-xylylene polymer Parylene™, and biocompatible metal coating.

37. (Withdrawn) An implantable constriction device according to claim 36, wherein

the biocompatible metal coating is selected from the group consisting of gold, silver and

titanium.

38. (Withdrawn) An implantable constriction device according to claim 32, wherein

hard silicone constitutes said base material.

39. (Withdrawn) An implantable constriction device according to claim 32, wherein

said first layer of said base material forms an inflatable tubing, and said second layer covers

said base material within said tubing.

40. (Withdrawn) An implantable constriction device according to claim 1, wherein

said base material forms an inflatable tubing and said property improving means comprises a

liquid impermeable coating coated on said base material.

41. (Withdrawn) An implantable constriction device according to claim 40, wherein

said tubing has an external surface of said base material and an internal surface of said base

material defining the interior of said tubing, said coating being coated on said external

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surface and/or internal surface.

- 42. (Withdrawn) An implantable constriction device according to claim 40, wherein said coating is selected from the group consisting of a poly-para-xylylene polymer ParyleneTM and a biocompatible metal coating.
- 43. (Withdrawn) An implantable constriction device according to claim 42, wherein the biocompatible metal coating is selected from the group consisting of gold, silver and titanium.
- 44. (Withdrawn) An implantable constriction device according to claim 40, wherein hard silicone constitutes said base material.
- 45. (Withdrawn) An implantable constriction device according to claim 40, wherein said base material forms two coaxial tubular layers and said property improving means comprises a tubular intermediate layer of a viscoelastic material located between said coaxial tubular layers.
- 46. (Withdrawn) An implantable constriction device according to claim 44, wherein said viscoelastic material comprises silicone gel, cellulose gel or collagen gel.

- 47. (Withdrawn) An implantable constriction device according to claim 40, wherein said base material forms an outer tubular layer and an inner arcuate layer attached to said outer tubular layer, said outer and inner layers defining a curved space extending longitudinally along said tubing, and said property improving means comprises viscoelastic material filling said space.
- 48. (Withdrawn) An implantable constriction device according to claim 47, wherein said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel, and collagen gel.
- 49. (Withdrawn) An implantable constriction device according to claim 1, wherein said property improving means comprises gas contained in a multiplicity of cavities formed in said base material to improve the flexibility of said composite structure.
- 50. (Withdrawn) An implantable constriction device according to claim 49, wherein said cavities are defined by net structures of said base material.
- 51. (Withdrawn) An implantable constriction device according to claim 49, wherein Tetrafluoroethylene polymer Teflon™ constitutes said base material.
 - 52. (Withdrawn) An implantable constriction device according to claim 49, wherein

said composite structure forms an inflatable tubing.

53. (Withdrawn/Currently Amended) An implantable constriction device for

forming a restricted stoma opening in the stomach or esophagus of a patient, comprising an

elongate composite structure adapted to constrict the stomach or esophagus of the patient,

wherein the composite structure includes an elongate biocompatible self-supporting base

material having surfaces exposed to aggressive body cells, when the constriction device is

implanted in the patient, and a cell barrier coating coated on said surfaces to prevent body

cells from breaking down the base material.

54. (Withdrawn) An implantable constriction device according to claim 53, wherein

said barrier coating is selected from the group consisting of a poly-para-xylylene polymer

Parylene™ and a biocompatible metal coating.

55. (Withdrawn) An implantable constriction device according to claim 54, wherein

the biocompatible metal coating is selected from the group consisting of gold, silver and

titanium.

56. (Currently Amended) An implantable constriction device for forming a restricted

stoma opening in the stomach of a patient, comprising:

elongate constricting means for externally constricting the stomach of the patient,

means for making the constricting means self-supporting, the self supporting means surrounding-covering the constricting means,

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the constricting means to either enlarge or restrict the stoma opening,

wherein said physical property improving means improves the resistance to aggressive body cells.

the means for making the constricting means self-supporting comprising a layer of polyurethane and a layer of silicone.

57. (Cancelled)

- 58. (Previously Presented) The implantable constriction device according to claim 56, wherein said property improving means comprises at least one layerlayer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.
- 59. (Currently Amended) The implantable constriction device according to claim 58, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene

polymer, a poly-para-xylylene polymer, and a biocompatible metal layer.

60. (Currently Amended) The implantable constriction device according to claim 59, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.

61. (Withdrawn) An implantable constriction device according to claim 56, wherein said property improving means improves the flexibility of said constricting means.

62. (Withdrawn) An implantable constriction device according to claim 61, wherein said property improving means comprises a core of a viscoelastic material covered with said self-supporting base material.

63. (Withdrawn) An implantable constriction device according to claim 62, wherein hard silicone constitutes said self-supporting means.

64. (Withdrawn) An implantable constriction device according to claim 62, wherein said viscoelastic material is selected from the group consisting of silicone gel, cellulose gel, and collagen gel.

65. (Withdrawn) An implantable constriction device according to claim 61, wherein

said property improving means comprises gas contained in a multiplicity of cavities formed

in said self-supporting means to improve the flexibility of said constricting means.

66. (Withdrawn) An implantable constriction device according to claim 65, wherein

said cavities are defined by net structures of said self-supporting means.

67. (Withdrawn) An implantable constriction device according to claim 65, wherein

Tetrafluoroethylene polymer Teflon™ constitutes said self-supporting means.

68. (Withdrawn) An implantable constriction device according to claim 56, wherein

said property improving means improves the fatigue resistance of said constricting means.

69. (Withdrawn) An implantable constriction device according to claim 68, wherein

said self-supporting means forms a first layer and said property improving means comprises

a second layer applied on said first layer, said second layer being more fatigue resistant than

said first layer.

70. (Withdrawn) An implantable constriction device according to claim 69, wherein

said second layer covers said first layer of said self-supporting means along a side of said

elongate constricting means that is intended to contact the esophagus or stomach.

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- 71. (Withdrawn) An implantable constriction device according to claim 69, wherein said second layer comprises a polyurethane layer.
- 72. (Withdrawn) An implantable constriction device according to claim 56, wherein said property improving means improves the liquid impermeability of said constricting means.
- 73. (Withdrawn) An implantable constriction device according to claim 72, wherein said self-supporting means forms an inflatable tubing and said property improving means comprises a liquid impermeable coating coated on said self-supporting means.
- 74. (Withdrawn) An implantable constriction device according to claim 73, wherein said tubing has an external surface of said self-supporting means and an internal surface of said self-supporting means defining the interior of said tubing, said coating being coated on said external surface and/or internal surface.
- 75. (Withdrawn) An implantable constriction device according to claim 73, wherein said coating is selected from the group consisting of a poly-para-xylylene polymer Parylene™ and a biocompatible metal coating.
 - 76. (Withdrawn) An implantable constriction device according to claim 75, wherein

titanium.

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the biocompatible metal coating is selected from the group consisting of gold, silver and

77. (Withdrawn) An implantable constriction device according to claim 55, wherein

hard silicon constitutes said self-supporting means.

78. (Cancelled)

79. Cancelled

80. (Withdrawn/Currently Amended) An implantable constriction device for

forming a restricted stoma opening in the stomach or esophagus-of a patient, comprising:

an elongate composite structure adapted to constrict the stomach or esophagus of the

patient,

a base material of said composite structure making said composite structure self-

supporting, said base material forming a first layer, and

a second layer applied on said first layer, said second layer being more fatigue

resistant than said first layer.

81. (Withdrawn/Currently Amended) An implantable constriction device for

forming a restricted stoma opening in the stomach or esophagus of a patient, comprising:

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an elongate composite structure adapted to constrict the stomach or esophagus of the patient,

a liquid semi-permeable base material of said composite structure forming an inflatable tubing and making said composite structure self-supporting, and a liquid impermeable coating coated on said base material.

82. (Withdrawn/Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach or esophagus of a patient, comprising: an elongate composite structure adapted to constrict the stomach or esophagus of the patient, and

a base material of said composite structure making said composite structure selfsupporting, said base material forming a multiplicity of gas-containing cavities to improve the flexibility of said composite structure.

83. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach or esophagus of a patient, comprising:

elongate means for externally constricting the stomach or esophagus of the patient,
means for making the constricting means self-supporting, the self supporting means
surrounding covering the constricting means,

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self

supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

wherein said property improving means improves the anti-friction properties of said constricting means,

- 84. (Currently Amended) The implantable constriction device according to claim 83, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.
- 85. (Currently Amended) The implantable constriction device according to claim 84, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.
- 86. (Currently Amended) The implantable constriction device according to claim 85, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.

87. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

elongate means for externally constricting the stomach of the patient,

means for making the constricting means self-supporting, the self supporting means surrounding-covering the constricting means,

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

- 88. (Currently Amended) The implantable constriction device according to claim 87, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.
 - 89. (Currently Amended) The implantable constriction device according to claim 88,

wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

- 90. (Currently Amended) The implantable constriction device according to claim 89, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.
- 91. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

elongate means for externally constricting the stomach of the patient,

means for making the constricting means self-supporting, the self supporting means surrounding covering the constricting means,

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

wherein said property improving means improves the softness of said constricting means.

- 92. (Currently Amended) The implantable constriction device according to claim 91, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach
- 93. (Currently Amended) The implantable constriction device according to claim 92, wherein said layer or coating comprises a viscoelastic material.
- 94. (Previously Presented) The implantable constriction device according to claim93, wherein said viscoelastic material comprises silicone gel, cellulose gel or collagen gel.
- 95. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

elongate means for externally constricting the stomach of the patient,

means for making the constricting means self-supporting, the self supporting means surrounding covering the constricting means.

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma

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opening,

wherein said property improving means improves the strength of said constricting

means.

the means for making the constricting means self-supporting comprising a layer of

polyurethane and a layer of silicone.

96. (Currently Amended) The implantable constriction device according to claim 95,

wherein said property improving means comprises at least one layer or coating on said self-

supporting means at least along a side of said elongate constricting means that is intended to

contact the stomach.

97. (Currently Amended) The implantable constriction device according to claim 96,

wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene

polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

98. (Previously Presented) The implantable constriction device according to claim

97, wherein the biocompatible metal layer is selected from the group consisting of gold,

silver and titanium.

99. (Currently Amended) An implantable constriction device for forming a

restricted stoma opening in the stomach of a patient, comprising:

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elongate means for externally constricting the stomach of the patient,

means for making the constricting means self-supporting, the self supporting means surrounding covering the constricting means.

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

wherein said property improving means improves the fatigue resistance of said constricting means,

the elongate means comprising silicone, and the self supporting means comprising polyurethane.

- 100. (Currently Amended) The implantable constriction device according to claim 99, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.
- 101. (Currently Amended) The implantable constriction device according to claim 100, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer

or coating.

102. (Currently Amended) The implantable constriction device according to claim 101, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.

103. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

elongate means for externally constricting the stomach of the patient.

means for making the constricting means self-supporting, the self supporting means surrounding covering the constricting means,

means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

wherein said property improving means improves the fatigue resistance of said constricting means.

- 104. (Currently Amended) The implantable constriction device according to claim 103, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.
- 105. (Currently Amended) The implantable constriction device according to claim 104, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.
- 106. (Currently Amended) The implantable constriction device according to claim 105, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.
- 107. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:
 - elongate means for externally constricting the stomach of the patient,
- means for making the constricting means self-supporting, the self supporting means surrounding covering the constricting means,
- means for improving at least one physical property of said constricting means other than self-supporting properties, said property improving means being applied on the self

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supporting means, and

means for adjusting the composite structure to either enlarge or restrict the stoma opening,

wherein said property improving means improves the aggressive body fluid resistant properties of said constricting means,

the means for making the constricting means self-supporting comprising a layer of polyurethane and a layer of silicone.

108. (Currently Amended) The implantable constriction device according to claim 107, wherein said property improving means comprises at least one layer or coating on said self-supporting means at least along a side of said elongate constricting means that is intended to contact the stomach.

109. (Currently Amended) The implantable constriction device according to claim 108, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

110. (Currently Amended) The implantable constriction device according to claim 109, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.

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111. (Currently Amended) The implantable constriction device according to claim 1,

wherein said layer or coating of said property improving means comprises an aggressive

body cell barrier layer or coating.

112. (Currently Amended) The implantable constriction device according to claim

111, wherein said layer or coating of said property improving means is applied on said base

material at least along a side of said elongate composite structure that is intended to contact

the stomach.

113. (Currently Amended) The implantable constriction device according to claim

112, wherein said layer or coating of said property improving means comprises a coating on

said base material.

114. (Currently Amended) The implantable constriction device according to claim

112, wherein said layer or coating is selected from the group consisting of a

tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer

or coating.

115. (Currently Amended) The implantable constriction device according to claim

114, wherein the biocompatible metal layer or coating is selected from the group consisting

of gold, silver and titanium.

116. (Currently Amended) The implantable constriction device according to claim 1, wherein said layer or coating of said property improving means has better aggressive body fluid resistant properties than said base material.

117. (Currently Amended) The implantable constriction device according to claim 116, wherein said layer or coating of said property improving means is applied on said base material at least along a side of said elongate composite structure that is intended to contact the stomach.

118. (Currently Amended) The implantable constriction device according to claim 117, wherein said layer or coating of said property improving means comprises a coating on said base material.

119. (Currently Amended) The implantable constriction device according to claim 117, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

120. (Currently Amended) The implantable constriction device according to claim

- 119, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.
- 121. (Currently Amended) The implantable constriction device according to claim 1, wherein said layer or coating of said property improving means has better anti-friction properties than the base material.
- 122. (Currently Amended) The implantable constriction device according to claim 121, wherein said layer or coating of said property improving means is applied on said base material at least along a side of said elongate composite structure that is intended to contact the stomach.
- 123. (Currently Amended) The implantable constriction device according to claim 122, wherein said layer or coating of said property improving means comprises a coating on said base material.
- 124. (Currently Amended) The implantable constriction device according to claim 122, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

- 125. (Currently Amended) The implantable constriction device according to claim 124, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.
- 126. (Currently Amended) The implantable constriction device according to claim 1, wherein said layer or coating of said property improving means comprises a liquid impermeable layer or coating.
- 127. (Currently Amended) The implantable constriction device according to claim 126, wherein said layer or coating of said property improving means is applied on said base material at least along a side of said elongate composite structure that is intended to contact the stomach.
- 128. (Currently Amended) The implantable constriction device according to claim 126, wherein said layer or coating of said property improving means comprises a coating on said base material.
- 129. (Currently Amended) The implantable constriction device according to claim 127, wherein said layer or coating is selected from the group consisting of a tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer or coating.

- 130. (Currently Amended) The implantable constriction device according to claim 129, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.
- 131. (Currently Amended) The implantable constriction device according to claim 1, wherein said layer or coating of said property improving means comprises a protective layer or coating softer than the base material.
- 132. (Currently Amended) The implantable constriction device according to claim 131, wherein said layer or coating of said property improving means is applied on said base material at least along a side of said elongate composite structure that is intended to contact the stomach.
- 133. (Currently Amended) The implantable constriction device according to claim 132, wherein said layer or coating of said property improving means comprises a viscoelastic material.
- 134. (Previously Presented) The implantable constriction device according to claim 132, wherein said viscoelastic material comprises silicone gel, cellulose gel or collagen gel.

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135. (Currently Amended) The implantable constriction device according to claim

134, wherein said silicone gel has a hardness less than 20 Shore.

136. (Currently Amended) The implantable constriction device according to claim 1,

wherein said layer or coating of said property improving means is stronger than the base

material.

137. (Currently Amended) The implantable constriction device according to claim

136, wherein said layer or coating of said property improving means is applied on said base

material at least along a side of said elongate composite structure that is intended to contact

the stomach-or esophagus.

138. (Currently Amended) The implantable constriction device according to claim

137, wherein said layer or coating of said property improving means comprises a coating on

said base material.

139. (Currently Amended) The implantable constriction device according to claim

137, wherein said layer or coating is selected from the group consisting of a

tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer

or coating.

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140. (Currently Amended) The implantable constriction device according to claim

139, wherein the biocompatible metal layer or coating is selected from the group consisting

of gold, silver and titanium.

141. (Currently Amended) The implantable constriction device according to claim 1,

wherein said layer or coating of said property improving means is more fatigue resistant than

the base material.

142. (Currently Amended) The implantable constriction device according to claim

141, wherein said layer or coating of said property improving means is applied on said base

material at least along a side of said elongate composite structure that is intended to contact

the stomach.

143. (Currently Amended) The implantable constriction device according to claim

142, wherein said layer or coating of said property improving means comprises a coating on

said base material.

144. (Currently Amended) The implantable constriction device according to claim

142, wherein said layer or coating is selected from the group consisting of a

tetrafluoroethylene polymer, a poly-para-xylylene polymer, and a biocompatible metal layer

or coating.

145. (Currently Amended) The implantable constriction device according to claim 144, wherein the biocompatible metal layer or coating is selected from the group consisting of gold, silver and titanium.

146. (Previously Presented) The implantable constriction device according to claim 1, wherein said base material is tubular.

147. (Currently Amended) The implantable constriction device according to claim 146, wherein said layer or coating of said property improving means is applied externally or internally on said tubular base material.

148. (Currently Amended) The implantable constriction device according to claim 146, wherein said property improving means comprises a first layer or coating applied externally on said tubular base material and a second layer or coating applied internally on said tubular base material.

149. (Currently Amended) The implantable constriction device according to claim 146, wherein said tubular base material comprises a double walled tubing having an external wall and an internal wall spaced from said external wall, whereby said external and internal walls define a space, and said layer or coating of said property improving means extends in

said space between said external and internal walls.

150. (Currently Amended) The implantable constriction device according to claim 149, wherein said layer or coating of said property improving means is softer than said external and internal walls of said base material.

151. (Currently Amended) The implantable constriction device according to claim 150, wherein said layer or coating of said property improving means comprises a viscoelastic material.

152. (Previously Presented) The implantable constriction device according to claim 151, wherein said composite structure comprises partition walls dividing said space between said external and internal walls into longitudinal cells, which are filled with said viscoelastic material.

153. (Previously Presented) The implantable constriction device according to claim 146, wherein said adjustment means comprises an elongate member sliding in said tubular base material.

154. (Previously Presented) The implantable constriction device according to claim 146, wherein said adjustment means is adapted to longitudinally displace said elongate FORSELL Application No. 10/623,801

composite structure to change the stoma opening.

155. (Currently Amended) An implantable constriction device for forming a restricted stoma opening in the stomach of a patient, comprising:

an elongate composite structure adapted to externally constrict the stomach, said elongate composite structure comprising:

an elongate member for constricting the stomach of the patient,

a base material attached to the elongate member, the base material making said composite structure self-supporting, and

property improving means for improving at least one physical property of said composite structure other than self-supporting properties, said property improving means including at least one layer or coating applied on said base material, and

an adjustment means to adjust the elongate member, and thereby, the composite structure to either enlarge or restrict the stoma opening.

the base material comprising a layer of polyurethane and a layer of silicone.